

# Tensi-Q

## Blood Pressure Monitor /

### Tensimeter Digital

No. Document: SPA-BM-PROD-199

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#### Foreword

Please read the User Manual carefully before using this product. The User Manual which describe the operating procedure should be followed strictly. This manual detailed introduce the steps must be noted when using the product, operation which may result in abnormal, the risk may cause personal injury and product damage and other contents, refer to the chapters for details. Any anomalies or personal injury and device damage arising from use, maintain, store do not follow requirements of the User Manual, our company is not responsible for the safety, reliability and performance guarantees! The manufacturer's warranty service does not cover such faults!

Our company has a factory record and user profile for each device, users enjoy free maintenance services for one year from the date or purchase. In order to facilitate us to provide you with a comprehensive and efficient maintenance service, please be sure to return the warranty card when you need repair service.

**Note: Please read the User Manual carefully before using this product.**

Described in this User manual is in accordance with practical situation of the product. In case of modifications and software upgrades, the information contained in this document is subject to change without notice.

#### The warning items

Before using this product, you should consider the safety and efficacy of the following described:

- Described each measurement results combined with clinical symptoms by qualified doctors.
- The reliability and operation of using this product whether meets the operation of this manual relate to the maintenance instructions.
- The intended operator of this product may be the patient.
- Do not perform maintenance and service while the device is in use.

**Warning: Replace accessories which not provided by our company may lead to the occurrence of errors. Replace adapters, cuffs at will may result in wrong measurement results. Without our company or other approved maintenance organizations trained service personnel should not try to maintain the product.**

#### Responsibility of operator:

- The operator must carefully read the User Manual before use this product, and strictly follow the operating procedure of the User Manual.
- Fully consider the security requirements during design, but the operator should not ignore the observation for the patient and the state of machines.
- The operator has the responsibility to provide the use condition of the product to our company.

#### Responsibility for our company

- Our company have the responsibility to provide qualified product which conform to company standard of this product.
- Our company will provide the circuit diagram, calibration method and other information at the request of the user to help the appropriate and qualified technicians to repair those parts designated by our company.
- Our company have the responsibility to complete product maintenance according to the contract.
- Our company have the responsibility to respond the requirements of user in time.
- In the following case, our company is responsible for the impact on the safety, reliability and performance of the device;

Assembly, addition, debugging, modification or repair are carried out by personnel approved by our company. The electrical facilities in the room are in compliance with the relevant requirement and the device is used in accordance with the User Manual.

**The User Manual is written by our company. All rights reserved.**

## CHAPTER1 FUNCTION AND PURPOSE

### 1.1 Main Functions

- Measure blood pressure and store the measurement results.
- Data storage function, up to 199 records can be stored.
- With data review interface which is convenient for reviewing blood pressure parameter.
- The screen will prompt message when the power is low.
- When the measurement result can not be obtained due to some factors during the measurement, the device will display the corresponding error information.
- Measurement units: mmHg or kPa, which can be switched by the button.
- With automatic shutdown function, if there is no operation, the device will automatically turn off.

### 1.2 Purpose

The device applies to measure the non-invasive blood pressure of human. Record parameter value of blood pressure to provide the reference for the health care professional.

## CHAPTER2 SAFETY PRECAUTIONS:

In order to use it correctly, please read the "Safety Precautions" carefully before using it.

Operators do not need professional training, but should use this product after fully understanding the requirements in this manual.

To prevent users from suffering damage or loss due to improper use, please refer to "Safety Precautions" and use this product properly.

**For safety reasons, be sure to comply with safety precautions.**

**Note: If not use correctly, it exists the possibility of damage for personnel and goods.**

Good damage means the damage of house, property, domestic animal and pet.

#### Contraindication: no.

#### Warning:

- You must not perform NIBP measurements on patients with sickle-cell disease or under any condition which the skin is damaged or expected to be damage.
- For patient with severe disturbances of blood coagulation, whether automatically measure the blood pressure should be based on the clinical evaluation, because limb friction with the cuff may cause the risk of hematoma.
- For severe blood circulation disorder or arrhythmia patients, please use the device under guidance of a doctor. If the arm is squeezed during measurement, it may cause acute internal hemorrhage or inaccurate measurement result.

#### Measurement Limitations:

To different patient conditions, the oscillometric measurement has certain limitation. The measurement is in search of regular arterial pressure pulse. In those circumstances when the patient's condition makes it difficult to detect, the measurement becomes unreliable and measuring time increases. The user should be aware that the following conditions could interfere with the measurement, making the measurement unreliable or longer to derive. In some cases, the patient's condition will make a measurement impossible.

#### Patient Movement

Measurements will be unreliable or can not perform if the patient is moving, shivering or having convulsions. These motions may interfere with the detection of the arterial pressure pulses. In addition, the measurement time will be prolonged.

#### Cardiac Arrhythmia's

Measurement will be unreliable and may not be possible if the patient's cardiac arrhythmia has caused an irregular heartbeat. The measuring time thus will be prolonged.

#### Heart-lung Machine

Measurements will not be possible if the patient is connected to a heart-lung machine.

#### Pressure Charges

Measurements will be unreliable and may not be possible if the patient's blood pressure is changing rapidly over the period of time during which the arterial pressure pulses are being analyzed to obtain the measurement.



#### Severe Shock

If the patient is in severe shock or hypothermia, measurements will be unreliable since reduced blood flow to the peripheries will cause reduced pulsation of the arteries.

#### Heart Rate Extremes

Measurements can not be made at a heart rate of less than 40 bpm and greater than 240 bpm.

#### Round Patient

The thick fat layer of body will reduce the measurement accuracy, because the fat come from the shock of arteries can not access the cuffs due to the damping.

#### WARNING:

**Self-diagnosis and treatment using measured results may be dangerous. Follow the instructions of your physician.**

Please hand measurement results to the doctor who knows your health and accept diagnosis.

**For Infant and the person who can't express oneself, please use the device under the guidance of a doctor.**

Otherwise it may cause accident or dissection.

**Please do not use for any other purpose except BP measurement.**

Otherwise it may cause accident or holdback.

**Please use special cuff.**

Otherwise it is possible that measurement result is incorrect.

**Please do not keep the cuff in the over-inflated state for a long time.**

Otherwise it may cause risk.

**Do not use the device in the case of there are flammable anesthetic gasses mixing with the air or nitrous oxide.**

Otherwise it may cause risk.

**If liquid splashes on the device or accessories, especially when liquids may enter the pipe or device, stop using and contact the service department.**

Otherwise it may cause risk.

**Dispose of the packaging material, observing the applicable waste control regulations and keeping it out of children's reach.**

Otherwise it may cause harm to the environment or children.

**Please use approved accessories for the device and check that the device and accessories are working properly and safely before use.**

Otherwise the measurement result may be inaccurate or an accident may occur.

**When the device is accidentally damp, it should be placed in a dry and ventilated place for a period of time to dissipate moisture.**

Otherwise the device may be damaged due to moisture.

**Do not store and transport the device outside the specified environment.**

Otherwise it may cause measurement error.

**It is recommended that you check if there is any damage on the device or the accessories regularly, if you find any damage, stop using it, and contact the biomedical engineer of the hospital or our Customer Service immediately. Do not disassemble, repair and modify the device without permission.**

Otherwise it cannot be accurately measured.

**This device can not be used on mobile transport platforms.**

Otherwise it may cause measurement error.

**This device can not be used on a tilted tabletop.**

Otherwise there is a risk of falling.

**Dispose of packaging materials, waste batteries and end-of-life products in accordance with local laws and regulations. The end-of-life products and materials are properly disposed of by the user in accordance with the authority's decree.**

Replace accessories which not provided by our company may lead to the occurrence of errors.

**Without our company or other approved maintenance organizations trained service personnel should not try to maintain the product.**

This device can only be used for one test object at a time.

**If the small parts on the device are inhaled or swallowed, please consult a doctor promptly.**

The device and accessories are processed with allergenic materials. If you are allergic to it, stop using this product.

**After pressing the power button, if the device has display fault such as white screen, blurred screen or no display content, please contact our company.**

The device shall comply with the standard IEC 80601-2-30: Particular requirements for basic safety and essential performance of automated non-invasive sphygmomanometers.

### 2.1 Operation for Power Adapter (Separate Sale)

**NOTE:**

The device can be powered by a power adapter that is a part of the medical electrical system. Be sure to use the dedicated medical grade power adapter of this device.

Otherwise it may cause trouble.

**Dedicated power adapter must use AC 100 V-240 V**

Otherwise it may cause fire or electric shock.

**When there is breakage of dedicated power adapter plug or wire, please do not use it.**

Otherwise it may cause fire or electric shock.

**Please do not plug or unplug the adapter on the socket with wet hands.**

Otherwise it may cause electric shock or injury.

**When using the power adapter to connect with the power socket, make sure the power socket is conveniently accessible, in order to timely disconnect from the power when emergency.**

### 2.2 Operation for Battery

**Please use 4 "AA" size manganese or alkaline batteries, do not use batteries of other types.**

Otherwise it may cause fire.

**Do not mix old and new batteries and batteries of different types**

Otherwise it may cause battery leakage, heat, rupture, and damage to Electronic Sphygmomanometer.

**Please don't put wrong the positive and negative of battery. When the batteries power exhausts, replace with four new batteries at the same time.**

**Please take out the batteries when you do not use the device for a long time (3 months or more).**

Otherwise it may cause battery leakage, heat, rupture, and damage to Electronic Sphygmomanometer.

**If electrolyte of the batteries immodestly gets in your eyes, immediately rinse with plenty of clean water.**

It will cause blindness or other hazards, should immediately go to the nearest hospital for treatment.

**If electrolyte of the batteries immodestly glues on the skin or the clothes, immediately rinse with plenty of clean water.**

Otherwise it may hurt the skin.

#### Advice

Do not strike or drop the device;

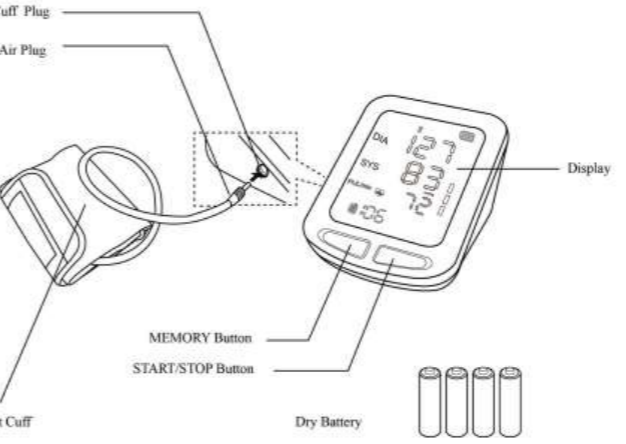
Do not inflate before the cuff wraps around the arm;

Do not infect the cuff and the air tube forcibly.

The device applies measurement Blood Pressure (BP) and Pulse of adult.

## CHAPTER3 MAIN UNIT

All products are in the box. Open box and confirm whether the product is whole.



All the operations to the Electronic Sphygmomanometer are through buttons. The names of the buttons are above them. They are:

- Left button is "M" button, under "OFF" state, press this button to enter the review interface (refer to Chapter 8 for details.).
- Right button is "START/STOP" button, under "OFF" state, press this button to enter measurement mode, inflate the cuff to measure blood pressure, press this button again to turn off the device.

### 6.2 Parameter setting

Under "OFF" state, press "M" button and "START/STOP" button simultaneously for 5 s to enter the setting interface, the default unit in this interface is "mmHg", short press "M" button to switch the unit between "mmHg" and "kPa".

## CHAPTER7 THE USAGE METHOD OF SPHYGMOMANOMETER

### 7.1 Accurate Measurement Way

**Measurement in quiet and relaxing state.**

- Adopt a comfortable sitting position, use back and arms to support the body.
- Place your elbow on a table, the palm faces up and the body is relaxed.
- The cuff is level with your heart.
- Feet flat on the floor, and do not cross your legs.



#### ADVICE:

Try to measure your blood pressure at the same time each day with the same arm and the same pose for consistency.

The high and low location of cuff will cause changes in measurement results.

Do not touch the sphygmomanometer, cuff and windpipe during measure.

**Measurements should be taken in a quiet place and the body relax.**

Remain still 4-5 minutes before measurement.

Do not talk and movement during the measurement. Relax the body, do not let the muscle activity.

Wait 4-5 minutes between measurements.

Do not use precision instrument near the Sphygmomanometer.

#### WARNING:

When repeatedly measuring, the accurate blood pressure value may not be measured due to congestion in the arm. Please measure after the blood flow is smooth.

Repeated measurement for a long period of time, limbs rubbing with the cuff may be accompanied by purpura, ischemia and nerve damage. When measurement a patient, it is necessary to frequently check the color, warmth and sensitivity of the distal of the limb. Once any abnormalities are observed, place the cuff

\*Self-diagnosis and treatment using measured results may be dangerous. Follow the instructions of your physician.

**NOTE:**

**Wait at least 4-5 minutes between measurements.**

- When repeatedly measuring, the accurate blood pressure value may not be measured due to congestion in the arm. Please measure after the blood flow is smooth.
- When the screen displays Err, the measure can't be carried out correctly.
- The minimum value of the patient's physiological signal is the minimum limit that the device can measure. The device may obtain inaccurate measurement results when it is operated below the minimum amplitude or minimum value of the patient's physiological signal.

\*The device will automatically turn off after five minutes in which there is no operation to the device, even if you forget to turn the power off.

**CHAPTER8 MEMORY FUNCTION**

The device can store NIBP values automatically, display up to 199 set of measurement results.

If 199 set of measurement data have been stored in current device, when saving the 200th set of data, the earliest set of data will be overwritten. If no measurement values, the memory values can be not numerated.

Memory function can not be used during measuring.

When there are no measurement values, " --- " will display on the review interface.

**8.1. Review the Memory Value**

Under "OFF" state, press "M" button to display the average value of the latest three set of data, when the number of measurement data is less than three groups, it will supplement automatically. Continue to press "M" button in current interface to view all measurement records.

**8.2. Delete Memory Value**

- Users can delete all memory values of the current user instead of separately delete one memory value.
- Under the memory interface, press "M" button and "START/STOP" button simultaneously for more than 5 s, after "DEL" appears on the screen, all memory values will be deleted.

**CAUTION:**

When querying the measurement records, please press "M" button continuously to query one by one.

**CHAPTER9 KEY AND SYMBOLS**

Your device may not contain all the following symbols.

Signal	Description	Signal	Description
	Attention! Please refer to the accompanying document (the user manual).		Attention! Please refer to the accompanying document (the user manual).
SYS	Systolic pressure	DIA	Diastolic pressure
MAP	Mean blood pressure	PUL	Pulse rate (bpm)
IPXX	Enclosure protection grade	EMC	Electromagnetic compatibility
	Recyclable	P/N	Material code of manufacturer
	Batch code		Use by date
	This way up		Fragile, handle with care
	Keep dry		Storage atmospheric pressure limitation
	Storage temperature limitation		Storage humidity limitation
	Manufacturer		Date of manufacture
	Batteries Power		Pulse rate (bpm)
	Flatting		Deflating
	Waste disposal mark, this symbol indicates that the waste of electrical and electronic equipment can not be disposed as an unclassified municipal waste and must be recovered separately.		Type BF applied parts
SN	Serial number		This item is compliant with Medical Device Directive 93/42/EEC of June 14, 1993, a directive of the European Economic Community.
	European Representative		Class II equipment
	Socket for Power Adapter		Interface for connecting cuff

**CHAPTER10 ERROR MESSAGE**

When the high pressure position appears "Err" and the low pressure position appears the error number, the measurement is not normal. (Error number are 02,04,06-16,19)

Error Mark	Causes	Solution
Err02 Err15	Function abnormal	Please contact us
Err04	Low battery	Please replace the battery or link adapter
Err06	The cuff is not wrapped correctly.	Wrap the cuff correctly (refer to Chapter 10)
Err07	Cuff plug fall off	Make sure the cuff plug is securely inserted in the windpipe (refer to Chapter 10)
Err08	Air pressure error	Keep arm, body still, measure again
Err09	The pulse signal is too weak or the cuff is loose.	Wrap the cuff correctly (refer to Chapter 10)
Err10	Cuff is blocked or squeezed	
Err14	Cuff leakage	Replace with a new cuff
Err11 Err12 Err13	The signal amplitude is too big owing to the arm or body moving or other reasons when measuring	
Err16 Err19	It takes too much time	Keep arm, body still, measure again

**CHAPTER11 TROUBLESHOOTING**

Abnormal Phenomenons	Causes	Solutions
BP measurement values too high or too low	Cuff is not connected correctly	Correctly connect cuff
	Talk or move arm in measurement	Keep quite and restart a measurement
	The turn up close oppress the	Take off the clothes, and restart a measurement
No Pressure	Cuff leakage	Buy new cuff

	The cuff windpipe is not correctly connected with cuff	Correctly connect
	Cuff not inflate	Contact us
Cuff deflate in short time	Loose cuff	Correctly tangle cuff
It can not carry on measurement, even if press the measurement		Return on the power and restart a measurement.
Abruptly turn the power off in adding pressure	No use for a long time, the batteries can be exhausted owing to the changed temperature	Replace all four batteries with new ones.
Hold the on/off button but can not start the device	Batteries are worn	Replace all four batteries with new ones.
	The batteries polarities is reversed	Check the battery installation for proper placement of the battery polarities
Cuff inflation start before press the measurement button		Stop using the device and contact us.
Cuff never deflation		Stop using the device and contact us.
Air pressure error	Deflation error	Pull out the cuff to deflate. Stop using the device and contact us.
	others	Keep arm, body still, measurement again
No press value displayed or the value unaltered when cuff inflating		Pull out the cuff to deflate. Stop using the device and contact us
Other phenomenon		Switch on the power once again and restart an operation.
		Replace the batteries.
		If no, please contact us.

Battery life	When the temperature is 23°C, limb circumference is 270 mm, the measured blood pressure is normal, 4 "AA" batteries can be used about 300 times.
Dimension	129(L)x101(W)x72(H) mm
Weight	281 gram (without batteries)
Safety classification	Class II equipment (power supplied by power adapter)/ Internally powered equipment (power supplied by batteries) Type BF applied part.
Service Life	The service life of the device is five years or 10000 times of BP measurement.
Date of manufacture	See the label
Accessories	Standard Configure: Adult Cuff: limb circumference 22-32 cm (upper arm center) User Manual, four "AA" alkaline batteries AC Adapter: Input: voltage: AC 100~240 V frequency: 50/60Hz rated current: AC 150 mA output: DC 5.0V±0.2V 1.0A

**APPENDIX**

**Guidance and manufacturer's declaration – electromagnetic emissions- for all EQUIPMENT and SYSTEMS**

**Guidance and manufacturer's declaration – electromagnetic emission**

The device is intended for use in the electromagnetic environment specified below. The customer or the user of the device should assure that it is used in such and environment.

Emission test	Compliance	Electromagnetic environment – guidance
RF emissions CISPR 11	Group 1	The device uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.
RF emission CISPR 11	Class B	The device is suitable for use in all establishments, including domestic establishments and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.
Harmonic emissions IEC61000-3-2	Class A	
Itage fluctuations/ flicker emissions IEC61000-3-3	Complies	

**Guidance and manufacturer's declaration – electromagnetic immunity-for all EQUIPMENT and SYSTEMS**

**Guidance and manufacturer's declaration – electromagnetic immunity**

The device is intended for use in the electromagnetic environment specified below. The customer or the user of the device should assure that it is used in such an environment.

Immunity test	IEC 60601 test level	Compliance level	Electromagnetic environment - guidance
Electrostatic discharge (ESD) IEC 61000-4-2	±6 kV contact ±8 kV air	±6 kV contact ±8 kV air	Floors should be wood, concrete or ceramic tile. If floor are covered with synthetic material, the relative humidity should be at least 30%.
Electrical fast transient/burst IEC 61000-4-4	±2 kV for power supply lines	±2kV for power supply lines	Mains power quality should be that of a typical commercial or hospital environment.
Surge IEC 61000-4-5	±1 kV differential mode ±2 kV common mode	±1 kV differential mode ±2 kV common mode	Mains power quality should be that of a typical commercial or hospital environment.
Voltage dips, short interruptions and voltage variations on power supply input lines IEC 61000-4-11	<5% Ur ( $\geq 95\%$ dip in Ur) for 0.5 cycle 40% Ur ( $\geq 60\%$ dip in Ur) for 5 cycles 70% Ur ( $\geq 30\%$ dip in Ur) for 25 cycles <5% Ur ( $\geq 95\%$ dip in Ur) for 5 sec	<5% Ur ( $\geq 95\%$ dip in Ur) for 0.5 cycle 40% Ur ( $\geq 60\%$ dip in Ur) for 5 cycles 70% Ur ( $\geq 30\%$ dip in Ur) for 25 cycles <5% Ur ( $\geq 95\%$ dip in Ur) for 5 sec	Mains power quality should be that of a typical commercial or hospital environment. The device can continue the operation during power mains interruptions due to the usage of battery.

NOTE: Ur is the a.c. mains voltage prior to application of the test level.

**Guidance and manufacturer's declaration – electromagnetic immunity –for EQUIPMENT and SYSTEMS that are not LIFE-SUPPORTING**

**Guidance and manufacturer's declaration – electromagnetic immunity**

The device is intended for use in the electromagnetic environment specified below. The customer or the user of the device should assure that it is used in such an environment.

Immunity test	IEC 60601 test level	Compliance level	Electromagnetic environment - guidance
Conducted RF IEC61000-4-6	3 Vrms 150 kHz to 80 MHz	3 Vrms	Portable and mobile RF communications equipment should be used no closer to any part of the device, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter.
Radiated RF IEC61000-4-3	3 V/m 80 MHz to 2.5 GHz	3 V/m	<b>Recommended separation distance</b> $d = \sqrt{\frac{3.5}{V_i} P}$

			$d = \sqrt{\frac{7}{E_i} P}$
			Where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and d is the recommended separation distance in metres (m). Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey, <sup>a</sup> should be less than the compliance level in each frequency range. <sup>b</sup> Interference may occur in the vicinity of equipment marked with the following symbol:

NOTE 1 At 80 MHz and 800 MHz, the higher frequency range applies.

NOTE 2 These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

A Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the device is used exceeds the applicable RF compliance level above, the device should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating the device.

B Over the frequency range 150 kHz to 80 MHz, field strengths should be less than 3 V/m.

Recommended separation distances between portable and mobile RF communications equipment and the device	Recommended separation distances between portable and mobile RF communications equipment and the EQUIPMENT or SYSTEM – for EQUIPMENT or SYSTEM that are not LIFE-SUPPORTING		
150 kHz to 80 MHz	80 MHz to 800 MHz	800 MHz to 2.5 GHz	


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